

DEC 27 2010

510(K) SUMMARY

This summary of 510(k) safety and effectiveness information is being submitted in accordance with the requirements of SMDA 1990 and 21 CFR §807.92(c).

The assigned 510(k) number is: K103583.

1. Submitter:

Shenzhen Mindray Bio-medical Electronics Co., LTD
Mindray Building, Keji 12th Road South, Hi-tech Industrial Park, Nanshan,
Shenzhen, 518057, P. R. China

Contact Person:

Meng Xianjun
Shenzhen Mindray Bio-medical Electronics Co., LTD
Mindray Building, Keji 12th Road South, Hi-tech Industrial Park,
Nanshan, Shenzhen, 518057, P. R. China
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Date Prepared: Sept. 25, 2010

2. Device Name: DC-7 Diagnostic Ultrasound System

Classification

Regulatory Class: II
Review Category: Tier II
21 CFR 892.1550 Ultrasonic Pulsed Doppler Imaging System (90-IYN)
21 CFR 892.1560 Ultrasonic Pulsed Echo Imaging System (90-IYO)
21 CFR 892.1570 Diagnostic Ultrasound Transducer (90-ITX)

3. Predicate Device(s):

DC-7 Diagnostic Ultrasound System is substantially equivalent to the following devices. They have the same technological characteristics, are comparable in key safety and effectiveness features, and have the same intended uses and basic operating modes as the predicate device.

Predicate Device	Manufacturer	Model	510(k) Control Number
1	Mindray	DC-7	K101041
2	Mindray	DC-3	K091941
3	GE	Voluson E8	K061682
4	GE	VIVID 7	K060542
5	GE	LOGIQ5 Expert	K032974
6	Siemens	Acuson CV70	K050240
7	Siemens	Acuson Sequoia 512	K052410
8	GE	LOGIQ P5	K060993

4. Device Description:

The DC-7 Diagnostic Ultrasound System is a general purpose, portable, software controlled, ultrasound diagnostic system. Its function is to acquire and display ultrasound images in B-mode, M-mode, Color mode, Color M mode, PW mode, CW mode, Power/DirPower mode, TDI mode or the combined mode (i.e. B/M Mode). This system is a Track 3 device that employs an array of probes that include linear array probe, convex array probe, phased array probe and volume probe with a frequency range of approximately 1.8 MHz to 12.0 MHz.

This modification will provide users with 3 additional transducers, some additional optional features called STIC, Stress Echo, iPage, and etc. These modifications all lead to overall quality and image enhancement.

5. Intended Use:

The DC-7 Diagnostic Ultrasound System is applicable for adults, pregnant women, pediatric patients and neonates. It is intended for use in abdominal, cardiac, small organ (breast, testes, thyroid, etc.), peripheral vascular, fetal, transrectal, transvaginal, pediatric, transcranial (adult cephalic and neonatal cephalic), musculoskeletal (conventional and superficial), intraoperative, and urology exams.

6. Non-clinical Tests:

DC-7 Diagnostic Ultrasound System has been evaluated for acoustic output, biocompatibility, cleaning and disinfection effectiveness as well as thermal, electrical and mechanical safety, and has been found to conform with applicable medical safety standards. This device has been designed to meet the following standards: UD 2, UD 3, IEC 60601-1, IEC 60601-1-1, IEC 60601-1-2, IEC 60601-2-37 ,IEC 60601-1-4 and ISO 10993-1.

7. Technological Characteristics:

The DC-7 Diagnostic Ultrasound System has the same technological characteristics with the predicate devices. All systems transmit ultrasonic energy into patients, then perform

post processing of received echoes to generate on-screen display of anatomic structures and fluid flow within the body. All system allow for specialized measurements of structures and flow, and calculations.

7. Conclusion:

The DC-7 Diagnostic Ultrasound System has the same technological characteristics, key safety and effectiveness features, and has the same intended uses and basic operating modes as the predicate devices.

Intended uses and other key features are consistent with traditional clinical practices, FDA guidelines and established methods of patient examination. The design, development and quality process of the manufacturer confirms with 21 CFR 820, ISO9001 and ISO13485 quality systems. The device conforms to applicable medical device safety standards.

Therefore, the DC-7 Diagnostic Ultrasound System is substantially equivalent with respect to safety and effectiveness to devices currently cleared for market.



DEPARTMENT OF HEALTH & HUMAN SERVICES

Public Health Service

Food and Drug Administration
10903 New Hampshire Avenue
Silver Spring, MD 20993

Shenzhen Mindray Bio-Medical Electronics Co., Ltd.
Attn: Mr. Jeff D. Rongero
Senior Project Engineer
Underwriters Laboratories, Inc.
12 Laboratory Drive
Research Triangle Park, NC 27709

DEC 27 2010

Re: K103583

Trade/Device Name: DC-7 Diagnostic Ultrasound System
Regulation Number: 21 CFR 892.1550
Regulation Name: Ultrasonic pulsed doppler imaging system
Regulatory Class: II
Product Code: IYN, IYO, and ITX
Dated: December 3, 2010
Received: December 7, 2010

Dear Mr. Rongero:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and we have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration.

This determination of substantial equivalence applies to the following transducers intended for use with the DC-7 Diagnostic Ultrasound System, as described in your premarket notification:

Transducer Model Number

<u>3C5A</u>	<u>L12-4</u>	<u>P7-3</u>
<u>C5-2</u>	<u>L7-3</u>	<u>P12-4</u>
<u>V10-4</u>	<u>L11-4</u>	<u>7LT4</u>
<u>V10-4B</u>	<u>L14-6</u>	<u>DE10-3</u>
<u>6C2</u>	<u>2P2</u>	<u>6LB7</u>
<u>7L4A</u>	<u>4CD4</u>	<u>6LE7</u>
<u>7L5</u>	<u>P4-2</u>	<u>CB10-4</u>

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to such additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 895. In addition, FDA may publish further announcements concerning your device in the Federal Register.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

This letter will allow you to begin marketing your device as described in your premarket notification. The FDA finding of substantial equivalence of your device to a legally marketed predicate device results in a classification for your device and thus permits your device to proceed to market.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please go to <http://www.fda.gov/AboutFDA/CentersOffices/CDRH/CDRHOFFICES/ucm115809.htm> for the Center for Devices and Radiological Health's (CDRH's) Office of Compliance. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to <http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm> for the CDRH's Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

If you have any questions regarding the content of this letter, please contact Paul Hardy at (301) 796-6542.

Sincerely yours,



David G. Brown, Ph.D.
Acting Director
Division of Radiological Devices
Office of *In Vitro* Diagnostic Device
Evaluation and Safety
Center for Devices and Radiological Health

Enclosure(s)

K103583

Indications for Use

510(k) Number (if known):

DEC 27 2010

Device Name: DC-7 Diagnostic Ultrasound System

Indications For Use:

The DC-7 Diagnostic Ultrasound System is applicable for adults, pregnant women, pediatric patients and neonates. It is intended for use in abdominal, cardiac, small organ (breast, testes, thyroid, etc.), peripheral vascular, fetal, transrectal, transvaginal, pediatric, transcranial (adult cephalic and neonatal cephalic), musculoskeletal (conventional and superficial), intraoperative, and urology exams.

Prescription Use AND/OR Over-The-Counter
(Part 21 CFR 801 Subpart D) (21 CFR 807 Subp

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ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of In Vitro Diagnostic Devices (OIVD)


(Division Sign-Off)
Division of Radiological Devices
Office of In Vitro Diagnostic Device Evaluation and Safety

Page 1 of 1

510K K103583

Diagnostic Ultrasound Indications for Use Form

System X Transducer _____
 Model: DC-7
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PWD	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal	P	P	P		P	P	P	Note 1,2,3,4,6,7
Abdominal	P	P	P	P	P	P	P	Note 1,2,3,4,5,6,7
Intraoperative (specify)*	P	P	P		P	P	P	Note 1,2,4,6,7
Intraoperative (Neuro)								
Laparoscopic								
Pediatric	P	P	P	P	P	P	P	Note 1,2,4,5,6,7
Small organ(specify)**	P	P	P		P	P	P	Note 1,2,4,6,7
Neonatal Cephalic	P	P	P	P	P	P	P	Note 1,2,4,5,6,7
Adult Cephalic	P	P	P	P	P	P	P	Note 1,2,4,5,6,7
Trans-rectal	P	P	P		P	P	P	Note 1,2,3,4,6,7
Trans-vaginal	P	P	P		P	P	P	Note 1,2,3,4,6,7
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional	P	P	P		P	P	P	Note 1,2,4,6,7
Musculo-skeletal Superficial	P	P	P		P	P	P	Note 1,2,4,6,7
Intravascular								
Cardiac Adult	P	P	P	P	P	P	P	Note 1,2,5,6,7
Cardiac Pediatric	P	P	P	P	P	P	P	Note 1,2,5,6,7
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular	P	P	P		P	P	P	Note 1,2,4,6,7
Other (specify)***	N	N	N		N	N	N	Note 1,2,4,6,7

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3: 4D(Real-time 3D)

Note 4: iScape

Note 5: TDI

Note 6: Color M

Note 7: Biopsy Guidance

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Concurrence of CDRH, Office of Device Evaluation(ODE)

Prescription USE (Per 21 CFR 801.109)


 (Division Sign-Off)

 Division of Radiological Devices
 Office of In Vitro Diagnostic Device Evaluation and Safety
510K 15163583

008-2

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: _____ 3CSA _____
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PWD	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal	P	P	P		P	P	P	Note 1, 2, 4,6,7
Abdominal	P	P	P		P	P	P	Note 1, 2, 4,6,7
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric	P	P	P		P	P	P	Note 1, 2, 4,6,7
Small organ(specify)**								
Neonatal Cephalic								
Adult Cephalic								
Trans-rectal								
Trans-vaginal								
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional	P	P	P		P	P	P	Note 1, 2, 4,6,7
Musculo-skeletal Superficial								
Intravascular								
Cardiac Adult								
Cardiac Pediatric								
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular	P	P	P		P	P	P	Note 1, 2, 4,6,7
Other (specify)***								

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D.

Note 3: 4D(Real-time 3D).

Note 4: iScape

Note 5: TDI

Note 6: Color M

Note 7: Biopsy Guidance

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Prescription USE (Per 21 CFR 801.109)



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 Division of Radiological Devices
 Office of In Vitro Diagnostic Device Evaluation and Safety

510K K102583

008-3

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: _____ C5-2 _____
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PWD	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal	P	P	P		P	P	P	Note 1, 2, 4,6,7
Abdominal	P	P	P		P	P	P	Note 1, 2, 4,6,7
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric	P	P	P		P	P	P	Note 1, 2, 4,6,7
Small organ(specify)**								
Neonatal Cephalic								
Adult Cephalic								
Trans-rectal								
Trans-vaginal								
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional	P	P	P		P	P	P	Note 1, 2, 4,6,7
Musculo-skeletal Superficial								
Intravascular								
Cardiac Adult								
Cardiac Pediatric								
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular	P	P	P		P	P	P	Note 1, 2, 4,6,7
Other (specify)***								

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power +PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3:4D(Real-time 3D)

Note 4: iScape

Note 5: TDI

Note 6: Color M

Note 7: Biopsy Guidance

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Prescription USE (Per 21 CFR 801.109)



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 Office of In Vitro Diagnostic Device Evaluation and Safety
 510K K103583 008-4

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer X
 Model: V10-4
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PWD	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal	P	P	P		P	P	P	Note 1, 2, 4,6,7
Abdominal								
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric								
Small organ(specify)**								
Neonatal Cephalic								
Adult Cephalic								
Trans-rectal	P	P	P		P	P	P	Note 1, 2, 4,6,7
Trans-vaginal	P	P	P		P	P	P	Note 1, 2, 4,6,7
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional								
Musculo-skeletal Superficial								
Intravascular								
Cardiac Adult								
Cardiac Pediatric								
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular								
Other (specify)***	N	N	N		N	N	N	Note 1,2,4,6,7

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3: 4D(Real-time 3D)

Note 4: iScape

Note 5: TDI

Note 6: Color M

Note 7: Biopsy Guidance

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Prescription USE (Per 21 CFR 801.109)



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 Division of Radiological Devices
 Office of In Vitro Diagnostic Device Evaluation and Safety

510K K103583

008-5

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: _____ V10-4B _____
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PWD	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal	P	P	P		P	P	P	Note 1, 2, 4,6,7
Abdominal								
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric								
Small organ(specify)**								
Neonatal Cephalic								
Adult Cephalic								
Trans-rectal	P	P	P		P	P	P	Note 1, 2, 4,6,7
Trans-vaginal	P	P	P		P	P	P	Note 1, 2, 4,6,7
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional								
Musculo-skeletal Superficial								
Intravascular								
Cardiac Adult								
Cardiac Pediatric								
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular								
Other (specify)***	N	N	N		N	N	N	Note 1,2,4,6,7

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3:4D(Real-time 3D)

Note 4: iScape

Note5: TDI

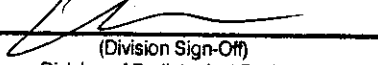
Note6: Color M

Note7: Biopsy Guidance

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Concurrence of CDRH, Office of Device Evaluation(ODE)

Prescription USE (Per 21 CFR 801.109)


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 Office of In Vitro Diagnostic Device Evaluation and Safety

510K K103583

Diagnostic Ultrasound Indications for Use Form

System Transducer X
 Model: 6C2
 510(k) Number(s)

Clinical Application	Mode of Operation							
	B	M	PWD	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal								
Abdominal	P	P	P		P	P	P	Note 1, 2, 4,6,7
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric	P	P	P		P	P	P	Note 1, 2, 4,6,7
Small organ(specify)**								
Neonatal Cephalic	P	P	P		P	P	P	Note 1, 2, 4,6,7
Adult Cephalic	P	P	P		P	P	P	Note 1, 2, 4,6,7
Trans-rectal								
Trans-vaginal								
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional	P	P	P		P	P	P	Note 1, 2, 4,6,7
Musculo-skeletal Superficial	P	P	P		P	P	P	Note 1, 2, 4,6,7
Intravascular								
Cardiac Adult								
Cardiac Pediatric								
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular	P	P	P		P	P	P	Note 1, 2, 4,6,7
Other (specify)***								

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3.4D(Real-time 3D)

Note 4: iScape

Note5: TDI

Note6: Color M

Note7: Biopsy Guidance:

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Concurrence of CDRH, Office of Device Evaluation(ODE)

Prescription USE (Per 21 CFR 801.109)


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 Division of Radiological Devices
 Office of In Vitro Diagnostic Device Evaluation and Safety

510K K03583

008-7

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: 7L4A
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PWD	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal								
Abdominal	P	P	P	P	P	P	P	Note 1,2,4,6,7
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric	P	P	P		P	P	P	Note 1,2,4,6,7
Small organ(specify)**	P	P	P		P	P	P	Note 1,2,4,6,7
Neonatal Cephalic	P	P	P		P	P	P	Note 1,2,4,6,7
Adult Cephalic								
Trans-rectal								
Trans-vaginal								
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional	P	P	P		P	P	P	Note 1,2,4,6,7
Musculo-skeletal Superficial	P	P	P		P	P	P	Note 1,2,4,6,7
Intravascular								
Cardiac Adult								
Cardiac Pediatric								
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular	P	P	P		P	P	P	Note 1,2,4,6,7
Other (specify)***								

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology, etc.

Note 1: Tissue Harmonic Imaging: The feature does not use contrast agents.

Note 2: Smart3D

Note 3:4D(Real-time 3D)

Note 4: iScape

Note5: TDI

Note6: Color M

Note7: Biopsy Guidance

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Concurrence of CDRH, Office of Device Evaluation(ODE)

Prescription USE (Per 21 CFR 801.109)


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 Division of Radiological Devices
 Office of In Vitro Diagnostic Device Evaluation and Safety

510K K103583

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: 7LS
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PWD	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal								
Abdominal	P	P	P		P	P	P	Note 1,2,4,6,7
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric	P	P	P		P	P	P	Note 1,2,4,6,7
Small organ(specify)**	P	P	P		P	P	P	Note 1,2,4,6,7
Neonatal Cephalic	P	P	P		P	P	P	Note 1,2,4,6,7
Adult Cephalic								
Trans-rectal								
Trans-vaginal								
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional	P	P	P		P	P	P	Note 1,2,4,6,7
Musculo-skeletal Superficial	P	P	P		P	P	P	Note 1,2,4,6,7
Intravascular								
Cardiac Adult								
Cardiac Pediatric								
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular	P	P	P		P	P	P	Note 1,2,4,6,7
Other (specify)***								

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3: 4D(Real-time 3D)

Note 4: iScape

Note5: TDI

Note6: Color M

Note7: Biopsy Guidance

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Concurrence of CDRH, Office of Device Evaluation(ODE)

Prescription USE (Per 21 CFR 801.109)



(Division Sign-Off)

Division of Radiological Devices
 Office of In Vitro Diagnostic Device Evaluation and Safety

510K h103583

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: L12-4
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PWD	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal								
Abdominal	P	P	P		P	P	P	Note 1,2,4,6,7
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric	P	P	P		P	P	P	Note 1,2,4,6,7
Small organ(specify)**	P	P	P		P	P	P	Note 1,2,4,6,7
Neonatal Cephalic	P	P	P		P	P	P	Note 1,2,4,6,7
Adult Cephalic								
Trans-rectal								
Trans-vaginal								
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional	P	P	P		P	P	P	Note 1,2,4,6,7
Musculo-skeletal Superficial	P	P	P		P	P	P	Note 1,2,4,6,7
Intravascular								
Cardiac Adult								
Cardiac Pediatric								
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular	P	P	P		P	P	P	Note 1,2,4,6,7
Other (specify)***								

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3: 4D(Real-time 3D)

Note 4: iScape

Note 5: TDI

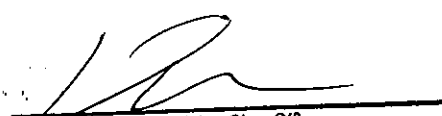
Note 6: Color M

Note 7: Biopsy Guidance

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Concurrence of CDRH, Office of Device Evaluation(ODE)

Prescription USE (Per 21 CFR 801.109)


 (Division Sign-Off)
 Division of Radiological Devices
 Office of In Vitro Diagnostic Device Evaluation and Safety

510K K103583

008-10

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: L7-3
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PWD	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal								
Abdominal	P	P	P		P	P	P	Note 1,2,4,6,7
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric	P	P	P		P	P	P	Note 1,2,4,6,7
Small organ(specify)**	P	P	P		P	P	P	Note 1,2,4,6,7
Neonatal Cephalic	P	P	P		P	P	P	Note 1,2,4,6,7
Adult Cephalic								
Trans-rectal								
Trans-vaginal								
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional	P	P	P		P	P	P	Note 1,2,4,6,7
Musculo-skeletal Superficial	P	P	P		P	P	P	Note 1,2,4,6,7
Intravascular								
Cardiac Adult								
Cardiac Pediatric								
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular	P	P	P		P	P	P	Note 1,2,4,6,7
Other (specify)***								

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3: 4D(Real-time 3D)

Note 4: iScape

Note5: TDI

Note6: Color M

Note7: Biopsy Guidance

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Concurrence of CDRH, Office of Device Evaluation(ODE)

Prescription USE (Per 21 CFR 801.109)


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 Division of Radiological Devices
 Office of In Vitro Diagnostic Device Evaluation and Safety

510K K103583

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: L11-4
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PWD	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal								
Abdominal	P	P	P		P	P	P	Note 1,2,4,6,7
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric	P	P	P		P	P	P	Note 1,2,4,6,7
Small organ(specify)**	P	P	P		P	P	P	Note 1,2,4,6,7
Neonatal Cephalic	P	P	P		P	P	P	Note 1,2,4,6,7
Adult Cephalic								
Trans-rectal								
Trans-vaginal								
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional	P	P	P		P	P	P	Note 1,2,4,6,7
Musculo-skeletal Superficial	P	P	P		P	P	P	Note 1,2,4,6,7
Intravascular								
Cardiac Adult								
Cardiac Pediatric								
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular	P	P	P		P	P	P	Note 1,2,4,6,7
Other (specify)***								

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3: 4D(Real-time 3D)

Note 4: iScape

Note5: TDI

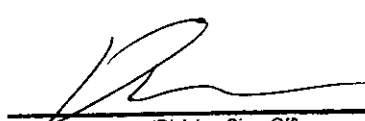
Note6: Color M

Note7: Biopsy Guidante

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Concurrence of CDRH, Office of Device Evaluation(ODE)

Prescription USE (Per 21 CFR 801.109)


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 Office of In Vitro Diagnostic Device Evaluation and Safety

510K 1103583

008-12

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: L14-6
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PWD	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal								
Abdominal	P	P	P		P	P	P	Note 1,2,4,6,7
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric	P	P	P		P	P	P	Note 1,2,4,6,7
Small organ(specify)**	P	P	P		P	P	P	Note 1,2,4,6,7
Neonatal Cephalic	P	P	P		P	P	P	Note 1,2,4,6,7
Adult Cephalic								
Trans-rectal								
Trans-vaginal								
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional	P	P	P		P	P	P	Note 1,2,4,6,7
Musculo-skeletal Superficial	P	P	P		P	P	P	Note 1,2,4,6,7
Intravascular								
Cardiac Adult								
Cardiac Pediatric								
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular	P	P	P		P		P	Note 1,2,4,6,7
Other (specify)***								

*new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3: 4D(Real-time 3D)

Note 4: iScape

Note 5: TDI

Note 6: Color M

Note 7: Biopsy Guidance

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Concurrence of CDRH, Office of Device Evaluation(ODE)

Prescription USE (Per 21 CFR 801.109)



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Division of Radiological Devices
 Office of In Vitro Diagnostic Device Evaluation and Safety

510K K103583

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: 2P2
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PW D	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal								
Abdominal	P	P	P	P	P	P	P	Note 1, 2,5,6,7
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric	P	P	P	P	P	P	P	Note 1, 2,5,6,7
Small organ(specify)**								
Neonatal Cephalic	P	P	P	P	P	P	P	Note 1, 2,5,6,7
Adult Cephalic	P	P	P	P	P	P	P	Note 1, 2,5,6,7
Trans-rectal								
Trans-vaginal								
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional								
Musculo-skeletal Superficial								
Intravascular								
Cardiac Adult	P	P	P	P	P	P	P	Note 1, 2,5,6,7
Cardiac Pediatric	P	P	P	P	P	P	P	Note 1, 2,5,6,7
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular								
Other (specify)***								

N=new indication, P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3:4D(Real-time 3D)

Note 4: iScape

Note5: TDI

Note6: Color M

Note7: Biopsy Guidance

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Concurrence of CDRH, Office of Device Evaluation(ODE)

Prescription USE (Per 21 CFR 801.109)

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Division of Radiological Devices

Office of In Vitro Diagnostic Device Evaluation and Safety

510K K103583

008-14

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: 4CD4
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PW D	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal	P	P	P		P	P	P	Note 1,2, 3, 4,6
Abdominal	P	P	P		P	P	P	Note 1,2, 3, 4,6
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric	P	P	P		P	P	P	Note 1,2, 4,6
Small organ(specify)**								
Neonatal Cephalic								
Adult Cephalic								
Trans-rectal								
Trans-vaginal								
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional								
Musculo-skeletal Superficial								
Intravascular								
Cardiac Adult								
Cardiac Pediatric								
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular								
Other (specify)***								

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3:4D(Real-time 3D)

Note 4: iScape

Note 5: TDI

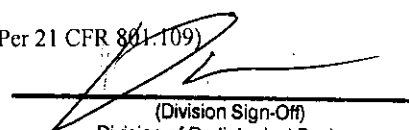
Note 6: Color M

Note 7: Biopsy Guidance

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 Office of In Vitro Diagnostic Device Evaluation and Safety

510K K103583

008-15

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: _____ P4-2
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PW D	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal								
Abdominal	P	P	P	P	P	P	P	Note 1, 2,5,6,7
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric	P	P	P	P	P	P	P	Note 1, 2,5,6,7
Small organ(specify)**								
Neonatal Cephalic	P	P	P	P	P	P	P	Note 1, 2,5,6,7
Adult Cephalic	P	P	P	P	P	P	P	Note 1, 2,5,6,7
Trans-rectal								
Trans-vaginal								
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional								
Musculo-skeletal Superficial								
Intravascular								
Cardiac Adult	P	P	P	P	P	P	P	Note 1, 2,5,6,7
Cardiac Pediatric	P	P	P	P	P	P	P	Note 1, 2,5,6,7
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular								
Other (specify)***								

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3:4D(Real-time 3D)

Note 4: iScape

Note 5: TDI

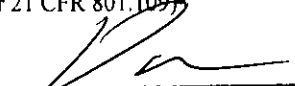
Note 6: Color M

Note 7: Biopsy Guidance

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Concurrence of CDRH, Office of Device Evaluation(ODE)

Prescription USE (Per 21 CFR 801.109)


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 Division of Radiological Devices
 Office of In Vitro Diagnostic Device Evaluation and Safety

510K K103583

008-16

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: _____ P7-3 _____
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PW D	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal								
Abdominal	P	P	P	P	P	P	P	Note 1, 2,5,6
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric	P	P	P	P	P	P	P	Note 1, 2,5,6
Small organ(specify)**								
Neonatal Cephalic	P	P	P	P	P	P	P	Note 1, 2,5,6
Adult Cephalic	P	P	P	P	P	P	P	Note 1, 2,5,6
Trans-rectal								
Trans-vaginal								
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional	P	P	P		P	P	P	Note 1, 2,6
Musculo-skeletal Superficial								
Intravascular								
Cardiac Adult	P	P	P	P	P	P	P	Note 1, 2,5,6
Cardiac Pediatric	P	P	P	P	P	P	P	Note 1, 2,5,6
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular								
Other (specify)***								

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3:4D(Real-time 3D)

Note 4: iScape

Note5: TDI

Note6: Color M

Note7: Biopsy Guidance

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510K K103583

008-17

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: P12-4
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PW D	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal								
Abdominal	P	P	P	P	P	P	P	Note 1, 2,5,6
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric	P	P	P	P	P	P	P	Note 1, 2,5,6
Small organ(specify)**								
Neonatal Cephalic	P	P	P	P	P	P	P	Note 1, 2,5,6
Adult Cephalic	P	P	P	P	P	P	P	Note 1, 2,5,6
Trans-rectal								
Trans-vaginal								
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional	P	P	P		P	P	P	Note 1, 2,6
Musculo-skeletal Superficial								
Intravascular								
Cardiac Adult	P	P	P	P	P	P	P	Note 1, 2,5,6
Cardiac Pediatric	P	P	P	P	P	P	P	Note 1, 2,5,6
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular								
Other (specify)***								

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3: 4D(Real-time 3D)

Note 4: iScape

Note 5: TDI

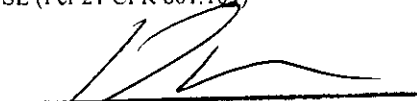
Note 6: Color M

Note 7: Biopsy Guidance

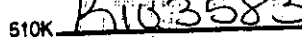
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Concurrence of CDRH, Office of Device Evaluation(ODE)

Prescription USE (Per 21 CFR 801.109)


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Office of In Vitro Diagnostic Device Evaluation and Safety

510K 

008-18

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: 7LT4
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PWD	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal								
Abdominal	P	P	P		P	P	P	Note 1,2,4,6,7
Intraoperative (specify)*	P	P	P		P	P	P	Note 1,2,4,6,7
Intraoperative (Neuro)								
Laparoscopic								
Pediatric	P	P	P		P	P	P	Note 1,2,4,6,7
Small organ(specify)**	P	P	P		P	P	P	Note 1,2,4,6,7
Neonatal Cephalic	P	P	P		P	P	P	Note 1,2,4,6,7
Adult Cephalic								
Trans-rectal								
Trans-vaginal								
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional	P	P	P		P	P	P	Note 1,2,4,6,7
Musculo-skeletal Superficial	P	P	P		P	P	P	Note 1,2,4,6,7
Intravascular								
Cardiac Adult								
Cardiac Pediatric								
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular	P	P	P		P	P	P	Note 1,2,4,6,7
Other (specify)***								

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3: 4D(Real-time 3D)

Note 4: iScape

Note 5: TDI

Note 6: Color M

Note 7: Biopsy Guidance

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Prescription USE (Per 21 CFR 801.109)


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 Division of Radiological Devices
 Office of In Vitro Diagnostic Device Evaluation and Safety

510K 1103583

008-19

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: DE10-3 _____
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PWD	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal	P	P	P		P	P	P	Note 1, 2,3, 4,6
Abdominal								
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric								
Small organ(specify)**								
Neonatal Cephalic								
Adult Cephalic								
Trans-rectal	P	P	P		P	P	P	Note 1, 2, 3,4,6
Trans-vaginal	P	P	P		P	P	P	Note 1, 2, 3,4,6
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional								
Musculo-skeletal Superficial								
Intravascular								
Cardiac Adult								
Cardiac Pediatric								
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular								
Other (specify)***								

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3:4D(Real-time 3D)

Note 4: iScape

Note 5: TDI

Note 6: Color M

Note 7: Biopsy Guidance

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Office of In Vitro Diagnostic Device Evaluation and Safety

008-20

510K K103583

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: 6LB7
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PWD	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal								
Abdominal								
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric								
Small organ(specify)**								
Neonatal Cephalic								
Adult Cephalic								
Trans-rectal	N	N	N		N	N	N	Note 1, 2, 4,6,7
Trans-vaginal								
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional								
Musculo-skeletal Superficial								
Intravascular								
Cardiac Adult								
Cardiac Pediatric								
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular								
Other (specify)***		N	N		N	N	N	Note 1,2,4,6,7

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3:4D(Real-time 3D)

Note 4: iScape

Note5: TDI

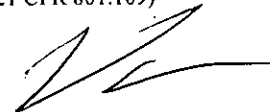
Note6: Color M

Note7: Biopsy Guidance

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Concurrence of CDRH, Office of Device Evaluation(ODE)

Prescription USE (Per 21 CFR 801.109)



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 Division of Radiological Devices
 Office of In Vitro Diagnostic Device Evaluation and Safety

610K 103583

008-21

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: 6LE7
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PWD	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal								
Abdominal								
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric								
Small organ(specify)**								
Neonatal Cephalic								
Adult Cephalic								
Trans-rectal	N	N	N		N	N	N	Note 1, 2, 4,6,7
Trans-vaginal								
Trans-urethral								
Trans-esoph.(non-Card.)								
Musculo-skeletal Conventional								
Musculo-skeletal Superficial								
Intravascular								
Cardiac Adult								
Cardiac Pediatric								
Intravascular (Cardiac)								
Trans-esoph.(Cardiac)								
Intra-Cardiac								
Peripheral Vascular								
Other (specify)***	N	N	N		N	N	N	Note 1,2,4,6,7

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3:4D(Real-time 3D)

Note 4: iScape

Note5: TDI

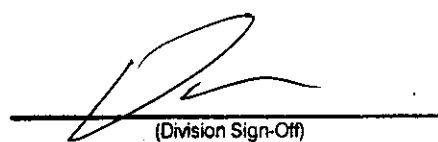
Note6: Color M

Note7: Biopsy Guidance

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Prescription USE (Per 21 CFR 801.109)



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 Office of In Vitro Diagnostic Device Evaluation and Safety

610K K1035823

008-22

Diagnostic Ultrasound Indications for Use Form

System _____ Transducer _____
 Model: CB10-4
 510(k) Number(s) _____

Clinical Application	Mode of Operation							
	B	M	PWD	CWD	Color Doppler	Amplitude Doppler	Combined (specify)	Other (specify)
Ophthalmic								
Fetal								
Abdominal								
Intraoperative (specify)*								
Intraoperative (Neuro)								
Laparoscopic								
Pediatric								
Small organ(specify)**								
Neonatal Cephalic								
Adult Cephalic								
Trans-rectal	N	N	N		N	N	N	Note 1, 2, 4,6,7
Trans-vaginal								
Trans-urethral								
Trans-esoph (non-Card.)								
Musculo-skeletal Conventional								
Musculo-skeletal Superficial								
Intravascular								
Cardiac Adult								
Cardiac Pediatric								
Intravascular (Cardiac)								
Trans-esoph (Cardiac)								
Intra-Cardiac								
Peripheral Vascular								
Other (specify)***	N	N	N		N	N	N	Note 1,2,4,6,7

N=new indication; P=previously cleared by FDA; E=added under Appendix E

Additional comments: Combined modes: B+M, PW+B, Color + B, Power + B, PW +Color+ B, Power + PW +B.

*Intraoperative includes abdominal, thoracic, and vascular etc.

**Small organ-breast, thyroid, testes, etc.

***Other use includes Urology.

Note 1: Tissue Harmonic Imaging. The feature does not use contrast agents.

Note 2: Smart3D

Note 3:4D(Real-time 3D)

Note 4: iScape

Note 5: TDI

Note 6: Color M

Note 7: Biopsy Guidance

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008-23